The FDA Safety Information and Adverse Event Reporting Program

**A. PATIENT INFORMATION**

<table>
<thead>
<tr>
<th>1. Patient Identifier</th>
<th>2. Age at Time of Event:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>In confidence</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>3. Sex</th>
<th>4. Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
<td>lbs</td>
</tr>
<tr>
<td>Male</td>
<td>kgs</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date of Birth:</th>
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</table>

**B. ADVERSE EVENT OR PRODUCT PROBLEM**

1. **Adverse Event or**

2. Outcomes Attributed to Adverse Event

3. **Date of Event** (mo/day/year)

4. **Date of This Report** (mo/day/year)

5. Describe Event or Problem

**C. SUSPECT MEDICATION(S)**

1. **Name** (Give labeled strength & mfr/labeler, if known)

2. **Dose**, **Frequency & Route Used**

3. **Therapy Dates** (If unknown, give duration) from/to (or best estimate)

4. **Diagnosis for Use** (Indication)

5. **Event Abated After Use Stopped or Dose Reduced?**

6. **Lot #** (If known)

7. **Exp. Date** (If known)

8. Event Reappeared After Reintroduction?

9. **NDC#** (For product problems only)

10. **Concomitant Medical Products and Therapy Dates** (Exclude treatment of event)

**D. SUSPECT MEDICAL DEVICE**

1. **Brand Name**

2. **Type of Device**

3. **Manufacturer Name, City and State**

4. **Model #**

5. **Operator of Device**

6. If Implanted, **Give Date** (mo/day/yr)

7. If Explanted, **Give Date** (mo/day/yr)

8. **Is this a Single-use Device that was Reprocessed and Reused on a Patient?**

9. **Is Yes to Item No. 8, Enter Name and Address of Reprocessor**

10. **Device Available for Evaluation?** (Do not send to FDA)

11. **Concomitant Medical Products and Therapy Dates** (Exclude treatment of event)

**E. INITIAL REPORTER**

1. **Name and Address**

2. **Health Professional?**

3. **Occupation**

4. **Initial Reporter Also Sent Report to FDA**

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For use by user-facilities, importers, distributors and manufacturers for MANDATORY reporting.

Submission of a report does not constitute an admission that medical personnel, user facility, importer, distributor, manufacturer or product caused or contributed to the event.

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Please type or use black ink.
### F. FOR USE BY USER FACILITY/IMPORTER (Devices Only)

1. Check One
   - User Facility
   - Importer

2. UF/Importer Report Number

3. User Facility or Importer Name/Address

4. Contact Person

5. Phone Number

6. Date User Facility or Importer Became Aware of Event (mo/day/yr)

7. Type of Report
   - Initial
   - Follow-up #

8. Date of This Report (mo/day/yr)

9. Approximate Age of Device

10. Event Problem Codes (Refer to coding manual)

11. Report Sent to FDA?
   - Yes (mo/day/yr)
   - No (mo/day/yr)

12. Location Where Event Occurred

13. Report Sent to Manufacturer?
   - Yes (mo/day/yr)
   - No (mo/day/yr)

14. Manufacturer Name/Address

### G. ALL MANUFACTURERS

1. Contact Office - Name/Address (and Manufacturing Site for Devices)

2. Phone Number

3. Report Source
   - Foreign
   - Study
   - Literature
   - Consumer
   - Health Professional
   - User Facility
   - Company Representative
   - Distributor
   - Other:

4. Date Received by Manufacturer (mo/day/yr)

5. (A/NDA #)

6. If IND, Give Protocol #
   - IND #
   - PLA #

7. Type of Report
   - Pre-1938
   - Yes
   - Other:

8. Adverse Event Term(s)
   - 5-day
   - 15-day
   - 10-day
   - Periodic
   - Initial
   - Follow-up #

### H. DEVICE MANUFACTURERS ONLY

1. Type of Reportable Event
   - Death
   - Serious Injury
   - Malfunction
   - Other:

2. If Follow-up, What Type?
   - Correction
   - Additional Information
   - Response to FDA Request
   - Device Evaluation

3. Device Evaluated by Manufacturer?
   - Not Returned to Manufacturer
   - Yes
   - Evaluation Summary Attached
   - No (Attach page to explain why not) or provide code:

4. Device Manufacture Date (mo/yr)

5. Labeled for Single Use?
   - Yes
   - No

6. Evaluation Codes (Refer to coding manual)
   - Method
   - Results
   - Conclusions

7. If Remedial Action Initiated, Check Type
   - Recall
   - Notification
   - Repair
   - Inspection
   - Replace
   - Patient Monitoring
   - Relabeling
   - Modification/Adjustment
   - Other:

8. Usage of Device
   - Initial Use of Device
   - Reuse
   - Unknown

9. If action reported to FDA under 21 USC 360i(f), list correction/removal reporting number:

10. Additional Manufacturer Narrative
   - and / or

11. Corrected Data

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The public reporting burden for this collection of information has been estimated to average one hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Department of Health and Human Services
Food and Drug Administration
MedWatch, HFD-410
5600 Fishers Lane
Rockville, MD 20857

Please DO NOT RETURN this form to this address.

OMB Statement:
"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number."